AMENDMENT

In the Claims:

The following listing of claims replaces all previous listings or versions thereof:

- 1. (Canceled)
- 2. (Original) The composition of claims 1, further defined as a A pharmaceutical composition comprising the at least one digitalis glycoside and an amorphous cyclodextrin, wherein the at least one digitalis glycoside includes oleandrin.
- 3. (Original) The composition of claim 2, wherein the pharmaceutical composition comprises one or more excipients.
- 4. (Original) The composition of claim 2, wherein pharmaceutical composition comprises one or more pharmaceutically acceptable antioxidants.
- 5. (Original) The composition of claim 2, wherein the pharmaceutical composition comprises one or more pharmaceutically acceptable preservatives.
- 6. (Original) The composition of claim 2, wherein the pharmaceutical composition comprises one or more pharmaceutically acceptable buffering agents.
- 7. (Original) The composition of claim 2, wherein the pharmaceutical composition comprises one or more pharmaceutically acceptable polysaccharides.
- 8. (Original) The composition of claim 3, wherein the said excipients comprises mannitol, sorbitol, fructose, glucose, lactose, sucrose, trehalose or any other water soluble sugar.
- 9. (Original) The composition of claim 4, wherein the said antioxidants comprise ascorbic acid, sodium ascorbate, sodium bisulfate, sodium metabisulfate, curcumin, curcumin derivatives, ursolic acid, resveratrol, resveratrol derivatives, alpha-lipoic acid or monothio glycerol.
- 10. (Original) The composition of claim 5, wherein the said preservatives comprise a methylparaben, methylparaben sodium, propylparaben, propylparaben sodium, benzalkonium chloride, or benzthonium chloride.

- 11. (Original) The composition of claim 6, wherein the said buffering agents comprise monobasic and dibasic sodium phosphate, sodium benzoate, potassium benzoate, sodium citrate, sodium acetate or sodium tartrate.
- 12. (Original) The composition of claim 7, wherein the polysaccharides comprise dextran sulfate, pectin, modified pectin, insoluble 1,3- β -D glucan, micronized 1,3- β -D glucan, soluble 1,3- β -D glucan, phosphorylated 1,3- β -D glucan, aminated 1,3- β -D glucan or carboxymethylated 1,3- β -D glucan, sulfated 1,3- β -D glucan.
- 13. (Currently amended) The composition of claim 1 claim 2, wherein the digitalis glycoside further comprises one or more of is oleandrin, neriifolin, odoroside A or H, ouabain (G-strophantin), cymarin, sarmentocymarin, periplocymarin, K-strophantin, thevetin A, cerberin, peruvoside, thevetosin, thevetin B, tanghinin, deacetyltanghinin, echujin, hongheloside G, honghelin, periplocin, strophantidol, nigrescin. uzarin, calotropin, cheiroside A, cheirotoxin, euonoside, euomonoside, lancetoxin A and B, kalanchoside, bryotoxin A-C, bryophyllin B, cotiledoside, tyledoside A-D, F and G, orbicuside A-C, alloglaucotoxin, corotoxin, coroglaucin, glaucorin, scillarene A and B, scilliroside, scilliglaucoside, scilliglaucosidin, scillirosidin, scillirubrosidin, scillirubroside, proscillaridin A, methyl-proscillaridin A, rubelin, convalloside, convallatoxin, bovoside A, glucobovoside A, bovoruboside, antiarin A, helleborin, hellebrin, adonidin, adonin, adonitoxin, thesiuside, digitoxin, gitoxin, gitalin, digoxin, F-gitonin, digitonin, lanatoside A-C, bufotalin, bufotalinin, bufotalidin, pseudobufotalin, acetyl-digitoxin, acetyl-oleandrin, beta-methyldigoxin or alphamethyldigoxin.
 - 14. 19. (Canceled)
- 20. (Original) The composition of claim 2 wherein said amorphous cyclodextrin has a degree of substitution of 2 to 7.
- 21. (Currently amended) The composition of elaim 1 claim 2 wherein the ratio by weight of digitalis glycoside to amorphous cyclodextrin is in the range of from 1:100 0.01 to 1 to 1.
- 22. (Currently amended) A process for preparing a pharmaceutical composition comprising admixing at least one digitalis glycoside, wherein one of said at least one digitalis

glycosides is oleandrin, with an amorphous cyclodextrin and rendering said composition pharmaceutically acceptable.

- 23. (Original) The process of claim 22, wherein the composition is rendered sterile by filtration.
- 24. (Original) The process of claim 22, wherein the composition is freeze-dried or lyophilized.
- 25. (Currently amended) A method of treating a cell proliferative disease in a subject comprising administering an amount of the composition of <u>claim 1 claim 2</u> that is effective to treat the cell proliferative disease, <u>wherein the proliferative disease is cancer</u>.
 - 26. (Original) The method of claim 25, wherein the subject is a human subject.
- 27. (Original) The method of claim 25, wherein the composition comprises the digitalis glycoside at a concentration of from 0.01 mg per mL to 10 mg per mL.
- 28. (Original) The method of claim 27, wherein the digitalis glycoside is at a concentration of from 0.04 mg per mL to 5 mg per mL.
- 29. (Original) The method of claim 25 wherein the composition is administered to the subject intramuscularly, intravenously or subcutaneously.
- 30. (Original) The method of claim 25, wherein the composition is administered orally, intranasally, rectally or vaginally.